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Citation for published version (APA):

Newbury, D. F., Gibson, J. L., Conti-Ramsden, G., Pickles, A., Durkin, K., & Toseeb, U. (Accepted/In press). Using Polygenic Profiles to Predict Variation in Language and Psychosocial Outcomes in Early and Middle Childhood. *Journal of Speech, Language, and Hearing Research*.

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Using Polygenic Profiles to Predict Variation in Language and Psychosocial Outcomes in Early and Middle Childhood

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Key Words: ALSPAC, Polygenic Risk, Language, Developmental Language Disorder, Social, Emotional, Conduct, Hyperactivity, Peer Problems, Prosocial, Psychopathology

Conflict of Interest: The authors have no conflicts of interest

Abstract

Purpose: Children with poor language tend to have worse psychosocial outcomes compared to their typically developing peers. The most common explanations for such adversities focus on developmental psychological processes whereby poor language triggers psychosocial difficulties. Here we investigate the possibility of shared biological effects by considering whether the same genetic variants which are thought to influence language development are also predictors of elevated psychosocial difficulties during childhood.

Method: Using data from the UK based Avon Longitudinal Study of Parents and Children (ALSPAC) we created a number of multi-SNP polygenic profile scores, based on language and reading candidate genes (*ATP2C2*, *CMIP*, *CNTNAP2*, *DCDC2*, *FOXP2*, & *KIAA0319*, 1229 SNPs) in a sample of 5,435 children.

Results: A polygenic profile score for expressive language (8 years) that was created in a discovery sample (n=2,718), predicted not only expressive language (8 years), but also peer problems (11 years) in a replication sample (n=2,717).

Conclusions: These findings provide a proof of concept for the use of such a polygenic approach in child language research when larger datasets become available. Our indicative findings suggest consideration should be given to concurrent intervention targeting both linguistic and psychosocial development as early language interventions may not stave off later psychosocial difficulties in children.

Using Polygenic Profiles to Predict Variation in Language and Psychosocial Outcomes in Early and Middle Childhood

There is a sound evidence for the heritable nature of language ability in children (Stromswold, 2001). The evidence concerning specific gene contributions is weaker: typically, single gene analyses account for only a very small proportion of trait variance or disorder (Simpson et al., 2015). Relationships between genetic variants within individuals, across populations and across genes means that the identification of a single variant is far from straightforward. This often means that the effects of variants do not replicate across studies or show differences in effect direction between cohorts. Within the current literature, it is hard to ascertain whether such variants represent false positive or heterogeneous effects (Nudel, Ceroni, Simpson, & Newbury, 2015). This state of affairs has led researchers interested in genetic effects to aggregate indicative findings concerning the relationships between specific alleles and particular traits into polygenic profile scores. The advantage of polygenic profiles is that they combine information from multiple variants and across genetic loci, enabling more comprehensive and powerful tests of the hypothesised ‘genetic burden’ associated with a condition (Levine et al., 2014; Mancuso et al., 2018; Wray et al., 2014).

The formation of polygenic profiles begins with a direct genetic association model within a large “discovery dataset”. This involves the investigation of correlations between individual genetic variants (usually single nucleotide polymorphisms or SNPs) and a behavioural outcome of interest. All SNPs that show association above a certain threshold (in terms of p-value or effect size) are then collapsed into a single weighted composite measure of genetic effects across multiple variants. Typically, scores are calculated across a range of thresholds in the discovery sample to allow the identification of the cut-off which confers maximal predictive power within the discovery sample set. Scores allow for the presence of false positive associations and datasets can be pruned for relationships between genetic variants (clumping). Once formulated, the polygenic profile score can be used within

a linear regression model to predict outcomes in a second independent “replication cohort”. This may be performed for any measure that is expected to show shared genetic effects with the behavioural measure of interest. The prediction accuracy is usually stated as an R^2 measure which represents the proportion of variance explained. Polygenic methods have proven particularly successful in disorders in which combinations of common variants predispose individuals to increased risk. Polygenic profiling has been used to indicate significant genetic overlaps between neuropsychiatric disorders (Ritter et al., 2017) as well as between cognitive ability and educational outcomes (Krapohl et al., 2018) and environmental influences and educational outcomes (Krapohl et al., 2017).

Such disorders are typically modelled under a “common disease-common variant” model. While genetic sequencing studies indicate that rare variants are also likely to play a role in neuropsychiatric disorders and traits, these will not be captured by polygenic profiling and are unlikely to be important in individual differences in language ability. In most instances, the overall predictive power of polygenic profiles remains low in terms of individual risk. Nonetheless, such profiles allow the identification of individuals at particularly high risk and can provide accurate indicators of the underlying risk interactions providing important information for risk modelling.

Given the lack of large-scale genome-wide associations for language-related measures, we selected robust candidate genes from the literature on genes on language and reading abilities and/or disorders, to construct targeted scores in the present study. Studies of targeted genes have indicated that some genes may have impacts across traits and disorders (Newbury et al., 2011; Scerri et al., 2011). Variants within the *KIAA0319* and *CNTNAP2* genes have been associated with both dyslexia and language impairment as well as reading and language measures in the general population (Newbury et al., 2011; Rice, Smith, & Gayan, 2009). In contrast, variants in *DCDC2* seem to be of particular relevance to reading

disability (Scerri et al., 2011) and those in *ATP2C2* to language impairment (Newbury et al., 2009). *FOXP2* appears to play a very specific role but may only be relevant in the presence of coding mutations with high-effect sizes (Morgan, Fisher, Scheffer, & Hildebrand, 2016). It has been suggested that variants within the *CMIP* gene may have alternative effects within different populations; the “risk” allele of rs12927866 has been associated with lower performance in tests of nonword repetition in language-impaired individuals but higher performance in the general population (Newbury et al., 2009). Such studies suggest that the role of risk variants may be modulated by the environment or genetic background of an individual and mirror findings of differential susceptibility in the psychiatry literature (Belsky, 1997).

In this study, we applied a polygenic approach in which we consider the effects of robust language and reading candidate genes within a single model. Specifically, we were interested in whether polygenic profiles, based on language and reading candidate genes (*ATP2C2*, *CMIP*, *CNTNAP2*, *DCDC2*, *FOXP2*, & *KIAA0319*), can be used to provide consistent predictors of *language outcomes* in early and middle childhood. This was assessed by the generation of polygenic profiles in a discovery cohort and the assessment of their correlation with the *same outcomes* in an independent replication cohort.

As well as predictors of outcomes, polygenic profiles can also be used to estimate the level of genetic overlap between contributing factors. Since genetic factors have previously been demonstrated to overlap between neurodevelopmental domains and across disorders (Anttila et al., 2018), we sought to investigate the possibility that polygenic profiles for language ability also contribute to the general development process. There is considerable support for this at the behavioural level where individuals with poor language, specifically those with Developmental Language Disorder (DLD), tend to have worse psychosocial outcomes compared to those without DLD (Beitchman et al., 2001; Botting, Durkin, Toseeb,

Pickles, & Conti-Ramsden, 2016; Botting, Toseeb, Pickles, Durkin, & Conti-Ramsden, 2016; Durkin, Toseeb, Botting, Pickles, & Conti-Ramsden, 2017; St Clair, Pickles, Durkin, & Conti-Ramsden, 2011; Yew & O'Kearney, 2015a, 2015b).

Such adverse outcomes are not inevitable. Whilst having DLD is certainly associated with risk of poorer psychosocial outcomes compared to unaffected individuals, these are not found invariably and the strength of any relationship may vary across different aspects of psychosocial functioning (Conti-Ramsden et al., 2018; Mok, Pickles, Durkin, & Conti-Ramsden, 2014; Pickles, Durkin, Mok, Toseeb, & Conti-Ramsden, 2016; Toseeb, Pickles, Durkin, Botting, & Conti-Ramsden, 2017). Investigating the aetiology of these differences in psychosocial outcomes in individuals with DLD speaks importantly to the debate about whether these problems have a common genetic origin or are linked in a developmental sequence. For example, in the latter explanation, a biologically-given language disorder could make social interaction difficult, and hence psychosocial difficulties (emotional instability, peer relationship problems, conduct disorder, hyperactivity, lack of prosociality) could follow developmentally (Durkin & Conti-Ramsden, 2007; Redmond & Rice, 1998).

While researchers have begun to address the complex task of uncovering genetic factors associated with language ability, scant attention has been paid to the question of whether the same genetic variants predict other characteristics in children with language disorders. In this study, we examined a model of shared genetic effects by considering whether polygenic profiles, based on language and reading candidate genes (*ATP2C2*, *CMIP*, *CNTNAP2*, *DCDC2*, *FOXP2*, & *KIAA0319*), can be used to provide consistent predictors of *psychosocial outcomes* in middle childhood.

Methods

Ethical Approval

Ethical approval for the study was obtained from the Avon Longitudinal Study of Parents and Children (ALSPAC) Ethics and Law Committee and the Local Research Ethics Committees. Ethical approval for the secondary analysis of existing ALSPAC data was obtained from the University of York Education Ethics Committee (reference: 18/5).

Study Sample

Data from the ALSPAC sample were used in this study. All pregnant women in the old administrative region of Avon, whose estimated delivery was between April 1991 to December 1992, were eligible to participate. The ALSPAC enrolled sample consisted of 14,775 live-born children from 15,247 pregnancies. This resulted in a total number of 15,458 children (including multiple births). Parents and children provided biological samples, questionnaire data, and took part in direct assessments. Full details of the cohort are reported elsewhere (Boyd et al., 2013; Fraser et al., 2013). Data used in this paper were mother reports on the child and direct assessments of the child by the research team at the following time points: the prenatal period (8 & 32 weeks gestation) and when the child was aged 15 months, 18 months, 24 months, 8 years, and 11 years old. Please note that the study website contains details of all the data that is available through a fully searchable data dictionary and variable search tool (<http://www.bristol.ac.uk/alspac/researchers/our-data/>).

An initial sample of 15,445 participants was provided by the ALSPAC study team. For the present study, the following exclusionary criteria were applied: children who did not have phenotypic data available from speech and language sessions at age 8 years old (n=8,062), children who scored below 60 or with incomplete data on performance IQ at age 8 years old (n=56), children who were born second in a multiple birth (twins or triplets) (n=100), unable to determine DLD status due to missing data or removed from DLD sample due to Autism Spectrum Disorder (ASD) and/or hearing loss (n=475), removed from no DLD sample due to ASD and/or hearing loss (n=391), missing phenotypic data (n=500), non-white

ethnicity (n=426). This resulted in a study sample of 5,435 (50% male), who were included in the genetic analyses.

Measures

Language measures. Eight measures of language development were collected. For the parent report measures at 15-24 months, a modified version of the MacArthur Communicative Development Inventory (CDI; Fenson et al., 1993) was used. The CDI has been shown to have good validity at a population level (Dale, Bates, Reznick, & Morisset, 1989; Feldman et al., 2005).

Vocabulary at 15 months. When the child was 15 months old, parents were given a list of 134 words/phrases and asked whether their child “understands but doesn’t say” (1), “understands and says” (2), or “neither” (0). Words/phrases were age-appropriate and included words such as “bed”, “nose” and “hot”. Responses were summed to create a score ranging from 0 to 268. Higher scores indicate better vocabulary.

Vocabulary at 24 months. When the child was 24 months old, a similar measure was used. Parents were given a list of 123 words/phrases and asked whether their child “understands” (1), “says” (2), or “neither” (0). Words were age-appropriate and included “hello”, “dinner”, and “chicken”. Responses were summed to create a score ranging from 0 to 246. Higher scores indicate higher vocabulary.

Receptive language at 15 months. Parents were shown a list of 12 phrases and asked whether their child understands. Samples phrases include “Are you sleepy?” and “Don’t touch”. Responses were coded on a binary scale (0=no, 1= yes) and then summed to create a score ranging from 0 to 12. Higher scores indicate better receptive language.

Grammar at 24 months. Parents were given four examples of grammar rules and asked whether their child has begun using these rules in their spoken language. Parents were asked about grammar rules such as adding “-ing” to the end of words and adding “-s” to

signify plural. Responses were coded as “not yet” (0), “sometimes” (1), “often” (2) and then summed to create a score ranging from 0 to 8. Higher scores indicate better grammar.

Receptive language at 8 years. The Weschler Objective Language Dimensions (WOLD, Rust, 1996) was used to measure receptive language. Only one of the two subsets were used in our study. The child was shown a picture and listened to a paragraph about the picture. The child then answered questions about what they had heard. The child was asked 16 questions. Responses were coded on a binary scale (0=incorrect, 1 = correct), yielding a summed score of between 0 and 16. Higher scores indicate better receptive language.

Expressive language at 8 years. The Weschler Objective Language Dimensions (WOLD, Rust, 1996) was used to measure expressive language. Only one of the two subsets was used. The child was shown ten pictures and asked to name them. Responses were coded on a binary scale (0= incorrect, 1= correct) and then summed to create a score ranging from 0 to 10. Higher scores indicate better expressive language.

Non-word repetition at 8 years. An adapted version of the Nonword Repetition Test (Gathercole et al, 1996) was used. The child was asked to listen and repeat out loud four each of 3, 4, and 5 syllable non-words. Responses were coded on a binary scale (0= incorrect, 1 = correct) and then summed to create a score ranging from 0 to 12. Higher scores indicate better non-word memory.

Pragmatic language at 9 years. The parent-report children’s communication checklist (CCC: Bishop, 1998) was used to measure pragmatic language when the child was 9 years old. A sum score was created using the sum of the inappropriate initiation, coherence, stereotyped conversation, use of conversational context, and conversational rapport subscales (ranging from 86 to 162). Higher scores indicate better pragmatic language.

Psychosocial measures. The parent report Strengths and Difficulties Questionnaire (SDQ: Goodman, 1997) was used to measure psychosocial outcomes at age 11 years. Sum scores were generated for each of the five subscales (emotional problems, peer problems, conduct problems, hyperactivity, & prosociality). The scores for each subscale ranged from 0 to 10, with higher scores indicating more difficulties for the problem sub-scales and higher prosociality.

Developmental Language Disorder Status (DLD)

Language measures provide an indication of individual differences in language ability but do not always reflect language difficulties which can be difficult to capture through measures in single domains. We therefore derived a measure of DLD using a previously reported framework (Scerri et al., 2011). Children were categorised as having DLD if they met at least two of the following four criteria: 1) pragmatic language >1 SD below standardised mean, 2) non-word repetition >1 SD below the standardised mean, 3) receptive language >1 SD below the standardised mean, or 4) positive response to “child has ever had speech/language therapy”. In line with recommendations regarding DLD diagnosis, children were excluded if they had a differentiating biomedical condition such as Autism Spectrum Disorder (ASD) or hearing problems (Bishop et al., 2017). ASD was defined as mothers responding positively that their child had Autism, Asperger’s, or Autism Spectrum Disorder at the age of 9 years. At age 7 years, children underwent a hearing test. Hearing problems were defined as bilateral hearing impairment, left unilateral hearing impairment, or right unilateral hearing impairment. Children with ASD or hearing loss were also excluded from the comparison sample. There were 346 children with DLD, which yielded a prevalence estimate of approximately 6%, which is consistent with prevalence data reported elsewhere (Norbury et al., 2016; Tomblin et al., 1997).

Our approach to identifying children with DLD, which involved selecting children at the tail end of the normal distribution on a number of language measures, allowed us to investigate DLD at a population level. Studies of clinical population may suffer from referral bias and may only represent children at the extreme end of the disorder. The sample of children with DLD that we identified will no doubt be a combination of those with a clinical diagnosis as well as those who are undiagnosed. Similar approaches have been previously used in other population-wide or community samples (see Forrest, Gibson, Halligan, & St Clair, 2018; Norbury et al., 2016; Reilly et al., 2010).

Descriptive statistics were used to generate profiles of the children with and without DLD. Psycholinguistic and psychosocial characteristics of these two groups, as shown in Table A1 in the appendices, support the use of such community-based approach. As a group, children with DLD had poorer early language ability (receptive language: 15 months, vocabulary: 15 & 24 months, and grammar: 24 months) compared to children without DLD. This was also the case later in childhood when they were aged 8-9 years. Children with DLD had lower levels of receptive, expressive, and pragmatic language as well as lower levels of non-word repetition compared to children without DLD. As a group, the children with DLD also had significantly more emotional problems, peer problems, conduct problems and had higher levels hyperactivity. They were also less prosocial compared to children without DLD.

In addition to this, descriptive statistics were run to investigate earlier language profiles of children with and without DLD. In total, 63% (n=209) of the children with DLD had impairment in receptive language and/or grammar aged 15-24 months compared to only 28% (n=1345) of children without DLD. These difficulties persisted at age 8-9 years when 79% (n=272) of children with DLD had impairment in receptive language and/or pragmatic

language compared to only 17% (n=864) of children without DLD. This supports the representativeness of this community-based sample of children with DLD.

Genetic Data

Quality control. Genetic data were obtained in a pre-processed format from the ALSPAC study team and included only SNP data for requested candidate genes. Participants were genotyped using the Illumina HumanHap550 quad chip genotyping platforms by 23andme subcontracting the Wellcome Trust Sanger Institute, Cambridge, UK and the Laboratory Corporation of America, Burlington, NC, US. Quality control of the data were performed prior to access. In short, individuals were excluded on the basis of gender mismatches; minimal or excessive heterozygosity; disproportionate levels of individual missingness (>3%) and insufficient sample replication ($IBD < 0.8$). Population stratification was assessed by multidimensional scaling analysis and compared with Hapmap II (release 22) European descent (CEU), Han Chinese, Japanese and Yoruba reference populations; all individuals with non-European ancestry were removed. SNPs with a minor allele frequency of < 1%, a call rate of < 95% or evidence for violations of Hardy-Weinberg equilibrium ($P < 5E-7$) were removed. Cryptic relatedness was measured as proportion of identity by descent ($IBD > 0.1$). Related participants that passed all other quality control thresholds were retained during subsequent phasing and imputation. 500,527 SNPs passed these quality control filters. This formed the pool for the selection of SNPs for the present study.

Selection of SNPs. In the absence of large genome-wide studies, we sought to increase the power of our polygenic approach through the *a priori* selection of candidate genes after a literature-based search. Genes that had shown previous robust evidence for association to language and/or reading within the ALSPAC population using common single nucleotide polymorphisms (SNPs) were included in the analyses (Newbury et al., 2009; Scerri et al., 2011). Although other genes have been previously associated with language,

these involved particular populations or single studies. The genes of interest and corresponding locations (hg38) were *ATP2C2* (chr16:84368615-84463732), *CMIP* (chr16:81445241-81709799), *CNTNAP2* (chr7:146116876-148415616), *DCDC2* (chr6:24174729-24357750), *FOXP2* (chr7:114426511-114693772), and *KIAA0319* (chr6:24541241-24645764). In total, there were 1,229 SNPs available at these locations. A summary of the number of SNPs by gene and location is shown in Table A2 in the appendices.

Splitting of dataset. The dataset was randomly split to generate two independent datasets for genetic analyses (referred to as the discovery and replication samples) using the “generate random” command in Stata/SE 14.2. This generated two approximately equal sized groups: discovery sample (n=2,718) and replication sample (n=2,717). There were no significant differences between these two sets of children on all measures of language and psychosocial outcomes ($ps > 0.05$).

Statistical Analyses

Association analysis. SNP data were analysed for allelic association within PLINK (Purcell et al., 2007) using linear regression models for the eight measures of language development. Eight measures were analysed across 1,229 SNPs. The association metrics for each SNP were then used to generate a best-fit polygenic profile for each of the dependent variables (language-measures) in the discovery cohort. The sensitivity of these profiles was evaluated by their ability to predict the same dependent variable within an independent replication set. This process is described in more detail below.

Polygenic analysis. Polygenic profile scores were calculated in the discovery cohort within the PRSice (v1.25) package (Euesden, Lewis, & O'Reilly, 2015). This package uses the effect sizes of individual SNPs (odds ratio or β) to estimate a weighted summation score that represents all variant effects within a single measure. Polygenic profiles were evaluated

across a range of P-value thresholds for the “base phenotype” in the “discovery” cohort. Best-fit scores at a given threshold were then used to predict the same base phenotype or an alternative “target” phenotype in an independent “replication” cohort. Polygenic profiles that significantly predict the outcome of the base measure in the replication set can be considered sensitive predictors of genetic risk. The overlaps between polygenic profiles and alternative target phenotypes can be considered to indicate the level of overlaps in genetic risk between base and target traits.

Polygenic profile scores were calculated at increments of 0.01 between P-value thresholds of 0 and 0.5 using the association results from the eight language measures in the discovery cohort. SNPs were thinned according to linkage disequilibrium and P-value within the discovery dataset using the “--clump” command in PRSice.

Each polygenic profile was tested within a step-wise procedure. First, the sensitivity of each of the generated polygenic profile was assessed by measuring the correlation of the score with the base language trait in the replication cohort. Secondly, the direct effects of the candidate genes were tested by exploring the ability of the polygenic profiles (independent variable) to predict language (DLD status) and psychosocial outcomes (SDQ subscales at age 11) as dependent variables in the replication cohort. Nominal p-values are presented in the text with Benjamini-Hochberg adjusted values also given in Table 3.

Results

Associations between phenotypes

Pairwise correlations were run between all language and psychosocial measures at all time points (see Table 1). Overall, there were positive correlations between language measures at similar time points. Therefore, for all parent report language measures that were taken when the children were 15-24 months old, there were significant correlations with medium to large effect sizes. Similarly, there were significant correlations between direct

measures of expressive and receptive language as well as non-word repetition taken when the children were 8 years old. The effect sizes for these were moderate. Pragmatic language when the children were 9 years old was also positively correlated with direct measures of language when the children were 8 years old, although the effect sizes were small. Analyses of language measures across different time points showed positive correlations but the effect sizes were generally small. Therefore, there was considerable variability between language development in early and middle childhood.

The findings of correlations between language and psychosocial measures were mixed. Higher language ability in early childhood was associated with fewer peer problems, lower levels of hyperactivity, and higher levels of prosociality at age 11 years old. On the other hand, language in early childhood was not associated with emotional and conduct problems when the children were 11 years old. Higher language ability in middle childhood was associated with fewer emotional (except for non-word repetition) and conduct problems as well as lower levels of hyperactivity. Therefore, we found some evidence for associations between language in early and middle childhood and psychosocial outcomes in middle childhood.

Genetic Associations

Polygenic profile scores, which incorporated the effects of common variants across six language-associated genes (*ATP2C2*, *CMIP*, *CNTNAP2*, *DCDC2*, *FOXP2*, and *KIAA0319*), were considered in relation to eight language development measures, DLD status and five SDQ subscales.

SNP based approach. Although association in the ALSPAC cohort has previously been described for some SNPs included in these analyses (Newbury et al., 2011; Scerri et al., 2011), the measures and sample set used in this study differ. We therefore first evaluated association between individual SNPs and the eight language measures to provide a baseline

of association prior to polygenic investigation. As shown in Table 2, at the single SNP level, no significant association was observed for any language measure tested in the discovery cohort after correction for multiple testing. Across all 9,832 tests performed (Hauser, Moutoussis, Dayan, & Dolan, 2017) fifteen had nominal P-values at the 10^{-4} level and the minimum nominal P-value was 1.77×10^{-4} . This was observed between the *CNTNAP2* variant rs9648690 and receptive language at 8 years of age (All SNPs with a p-value of 10^{-4} are shown in Appendix Table A3). In general, the language measures at 8 and 9 years of age showed a higher level of association than measures taken at 15 or 24 months of age. These data illustrate the difficulties associated with single SNP analyses across candidate genes and support the rationale for polygenic analyses, which collapse multiple genetic measures into a single factor and may be more robust to differences between sample sets.

[TABLE 2 HERE]

Consistency of polygenic profile scores. Polygenic profile scores were calculated for each of the eight language measures on the basis of the effect sizes for all SNPs investigated in the discovery cohort. These scores were then evaluated for their association with the same base measure in the independent replication sample under the same threshold allowing the identification of scores and measures that provide a sensitive marker of genetic effects across samples (Table 3). One of the eight polygenic profile scores generated (expressive language at 8 years of age) did provide a consistent marker for genetic effects upon language outcomes. The best-fit for this polygenic profile score was found at a P-threshold of 0.23 and explains 0.18% of trait variance (R^2) in the replication set (nominal $P=0.042$) (Figure 1). After clumping for linkage disequilibrium and P-value, this score was based upon 65 SNPs spread across all six candidate genes included. Seven of the SNPs fell in reading candidate genes (*DCDC2* and *KIAA0319*) with the remainder falling in language candidate genes (*ATP2C2*, *CMIP*, *CNTNAP2*, and *FOXP2*).

[FIGURE 1 HERE]

[TABLE 3 HERE]

Polygenic effects on other outcomes. The effects of the polygenic profile score upon expressive language at 8 years was found to be correlated with peer problems at 11 years of age suggesting genetic overlaps between expressive language and this outcome (Table 3). To inform our understanding of the extent of overlaps, we calculated the best-fit threshold across both outcome measures in the replication set. At a P-value threshold of 0.23 (as maximised in the discovery sample), the expressive language profile score explained 0.22% of variation in peer problems at age 11 years (nominal $P=0.049$). In contrast the best-fit for the prediction of peer problems at 11 years was found at a P-threshold of 0.06 and included 27 SNPs representing a subset of those contributing to expressive language. This score explained 0.43% of the trait variance in peer problems (R^2) in the replication set (nominal $P=0.0058$, Benjamini-Hochberg corrected p-value of 0.036 at a FDR of 0.05) (Table 3, Figure 2).

Discussion

Using polygenic profile scores, we investigated genetic effects upon language and psychosocial outcomes. Polygenic profile scores indicated some evidence of association across the SNPs tested, and the profile score for expressive language at 8 years provided a consistent marker across the candidate genes. Of particular interest, the expressive language profile score at 8 years was significantly associated with peer problems at age 11 years. These findings are consistent with behavioural models in which early language difficulties increase the risk of psychosocial difficulties.

The polygenic profile scores used here were based upon six genes that have previously been associated with language and/or reading. We hypothesised that this targeted approach would increase the likelihood of constructing meaningful polygenic profiles within the moderate sample sizes available (Dudbridge, 2013). In support of this hypothesis, we

found that variation across these candidate genes could provide a consistent marker of the genetic effects upon expressive language, although the proportion of variance explained remained low throughout ($<0.3\%$). All genes contributed to the risk model with 5% (65 of the 1229 SNPs tested) of variants contributing to the best-fit polygenic profile. Whilst the proportion of variance explained is low, this is not uncommon. In other fields, such as education, earlier studies of polygenic effects explained $\sim 2\%$ of variance (Rietveld et al., 2013). In the most recent work, larger datasets have allowed for the identification of more SNPs and so increasing the variance explained to $\sim 13\%$ (Lee et al., 2018). Similarly, larger sample sets and Genome Wide Association (GWAS) studies will allow the relative evaluation of the targeted loci studied in this paper. *A priori* candidate genes have been substantiated within larger polygenic studies (Ritchie et al., 2019), although many candidate genes do not replicate at the GWAS level (Border et al., 2019; Johnson et al., 2017). Despite the small proportion of variance explained, our findings support the role of common variants of small effect sizes within a complex genetic model of language development and reiterate the utility of polygenic profiles, which capture multiple effects within a single score.

Furthermore, polygenic profile scores of expressive language were able to predict peer problems at age 11 years indicating that genetic overlaps between this measure and psychosocial outcomes (Table 3). These findings support the previous behavioural findings which show that language ability is correlated with psychosocial outcomes (Conti-Ramsden & Botting, 2004; Forrest et al., 2018; Mok et al., 2014). In addition, our study extends these observations in that it suggests that expressive language difficulties may increase the concomitant peer problems through shared biological pathways informing our understanding of the routes to psychosocial difficulties in children with language disorder. If future research replicates our findings and is able to explain a larger proportion of variance, such a finding is important as it may indicate the need for concurrent intervention targeting both

linguistic and psychosocial development, rather than assuming early language interventions will stave off later emergence of peer problems.

We found that the best-fit P-value thresholds for the prediction of peer problems ($P_T=0.106$ respectively) formed a subset of those which provided the most consistent score for expressive language alone ($P_T=0.23$). This finding indicates that within the polygenic model different subsets of SNPs may be more relevant to different outcomes. Although the actual number of variants differed between thresholds, the weighting of individual SNPs was fixed between thresholds and the SNPs that contributed to peer problems formed a direct subset of those associated with expressive language. Specifically, we found that 42% (27 of 65) of the variants contributing to expressive language also contributed to peer problems at age 11 years. The relative numbers of variants from each gene was consistent across thresholds and, in general, these ratios reflected gene size. This therefore supports the relative role of all six candidate genes in language development.

The methods reported here provide a useful approach to the investigation of relationships between genetic risks and environmental factors. Through the application of polygenic profiles, we have demonstrated potential shared genetic relationships between expressive language and peer problems. These findings provide further evidence of the role of genes in language development and emphasise the importance of larger-scale studies to identify specific factors that moderate risk and mediate positive psychosocial outcomes for children affected by language disorder. Likewise, investigations of genetic comorbidities between language disorder and psychosocial outcomes represent an interesting future direction (Bulik-Sullivan et al., 2015). Application of this approach has revealed substantial overlaps between neurodevelopmental disorders and in relation to educational attainment (Gialluisi et al., 2019; Grove et al., 2019). Such studies allow biological insights into disease

mechanisms and may help to target intervention to individuals who will best benefit from additional support in terms of long-term outcomes.

An additional finding of note concerns the prosocial subscale of the SDQ. Previous evidence indicates that individuals with DLD show relatively stable prosociality scores from late childhood to 24 years (Lindsay & Dockrell, 2012; Toseeb et al., 2017). The finding that prosociality was within the normal range for both those with DLD and those without (Table A1), provide further support for the inference that prosociality is a characteristic of relative strength in young people with DLD measure (in contrast to their less favourable scores on the other SDQ subscales). We obtained no indication that prosociality scores could be predicted by polygenic profiles, suggesting that this aspect of psychosocial functioning may either be influenced by other genetic factors or be nurtured by socialization processes (or some interaction of these factors). We cannot resolve this issue on the basis of the present data, but the findings do add to an accumulating body of evidence that this strength in prosociality in those with DLD may be sufficiently robust to warrant incorporation in therapeutic work (Conti-Ramsden & Durkin, 2016; Toseeb et al., 2017).

When considering the findings of this study, some drawbacks should be borne in mind. This study considers only a small number of genes which, between them, account for only a small proportion of genetic risk. Thus, although the patterns that we observed support existing behavioural data, we must qualify these observations with the fact that these effects only represent a small proportion of genetic liability and a small part of the complex picture of risks relevant to language disorders. Nonetheless, in the absence of a large-scale picture of genome-wide effects in language disorder, the focus of this work upon a small set of robust candidate genes allowed the derivation of consistent polygenic profile of language disorder, albeit with a small effect size. A functional candidate approach, like the one we have reported here, has previously been shown to provide an accurate way of focusing polygenic profile

studies, for example in fibrogen pathways in cardiovascular disease (Cronjé et al., 2017). In this instance, the use of a specified set of candidate genes allowed the further exploration of relative subsets in relation to outcome measures. This study illustrates the utility of polygenic models in the study of language disorders and when larger sample become available, can be extended to a genome-wide model.

We should also note that only one of the P-values found in the current study survived multiple testing corrections. Although polygenic profiles allow the reduction of dimensions through the consideration of a single weighted-risk score, they still involve multiple tests, especially when high-resolution best-fit modelling is employed (Euesden et al., 2015). The use of hypothesis-driven, pre-defined models offsets this issue to some extent as we have some *a priori* expectation of the patterns we will observe from the behavioural literature. Nonetheless, these findings require extension to, and replication in, larger, independent datasets to claim significance.

Overall, our study illustrates the utility of polygenic methods in the study of children's language development. We found preliminary evidence that polygenic profiles for expressive language can be used to predict expressive language and peer problems in an independent sample. Our findings point to particular language and psychosocial outcomes that appear to be associated with genetic risk of language disorder.

Acknowledgements

We are extremely grateful to all the families who took part in this study, the midwives for their help in recruiting them, and the whole ALSPAC team, which includes interviewers, computer and laboratory technicians, clerical workers, research scientists, volunteers, managers, receptionists and nurses. We would also like to acknowledge the contributions of Witold Orlik and Peter Clough during the preliminary stages of the project.

The UK Medical Research Council and Wellcome (Grant ref: 102215/2/13/2) and the University of Bristol provide core support for ALSPAC. This publication is the work of the authors and Newbury, Gibson, Conti-Ramsden, Durkin, and Toseeb will serve as guarantors for the contents of this paper. A comprehensive list of grants funding is available on the ALSPAC website (<http://www.bristol.ac.uk/alspac/external/documents/grant-acknowledgements.pdf>); The analysis undertaken in this paper was specifically funded by Economic and Social Research Council (Grant ref: ES/P001955/1 & ES/P001955/2. G. CR is supported by the NIHR Manchester. AP is partially supported by NIHR NF-SI-0617-10120 and the Biomedical Research Centre at South London and Maudsley NHS Foundation Trust and King's College London. The views expressed are those of the author(s) and not necessarily those of the NHS, the NIHR or the Department of Health and Social Care.

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Table 1

Pairwise correlations between all phenotypes

	1.	2.	3.	4.	5.	6.	7.	8.	9.	10.	11.	12.	13.
1.Vocabulary 15months	1												
2.Vocabulary 24months	0.59***	1											
3.Receptive 15months	0.61***	0.41***	1										
4.Grammar 24months	0.47***	0.74***	0.28***	1									
5.Receptive language 8years	0.04*	0.11***	0.00	0.09***	1								
6.Expressive language 8years	0.10***	0.21***	0.03*	0.19***	0.37***	1							
7.Non-word repetition 8years	0.13***	0.30***	0.06***	0.26***	0.20***	0.30***	1						
8.Pragmatic language 9years	0.08***	0.17***	0.05***	0.14***	0.10***	0.15***	0.16***	1					
9.Emotional problems 11years	0.02	0.01	0.00	0.01	-0.06**	-0.03*	-0.02	-0.22***	1				
10.Peer problems 11years	-0.05**	-0.10***	-0.07***	-0.07***	0.01	0.00	-0.04**	-0.30***	0.35***	1			
11.Conduct problems 11years	-0.01	-0.04*	-0.03	-0.02	-0.06***	-0.08***	-0.07***	-0.30***	0.29***	0.25***	1		
12.Hyperactivity 11years	-0.11***	-0.15***	-0.07***	-0.12***	-0.08***	-0.12***	-0.11***	-0.44***	0.25***	0.23***	0.47***	1	
13.Prosociality 11years	0.09***	0.07***	0.13***	0.05***	-0.02	-0.02	-0.03*	0.16***	-0.10***	-0.20***	-0.41***	-0.32***	1

* p<.05, ** p<.01, *** p<.001.

Table 2

Genetic association to language measures in discovery cohort

Gene	Vocabulary 15months	Vocabulary 24months	Receptive language 15months	Grammar 24months	Receptive Language 8years	Expressive Language 8years	Nonword repetition 8years	Pragmatic language 9years	minP	minP trait
<i>ATP2C2</i>	0.01513	0.00543	0.003633	0.01061	0.000631	0.000662	0.01517	0.016	0.000631	Receptive language
<i>CMIP</i>	0.009171	0.01004	0.001037	0.05581	0.00034	0.000582	0.006129	0.01474	0.00034	Receptive language
<i>CNTNAP2</i>	0.003648	0.000648	0.002706	0.002598	0.000177	0.004907	0.000244	0.000922	0.000177	Receptive language
<i>DCDC2</i>	0.05209	0.004193	0.01358	0.04792	0.000295	0.008146	0.01634	0.03848	0.000295	Receptive language
<i>FOXP2</i>	0.09811	0.03679	0.03812	0.1279	0.06864	0.2249	0.2641	0.001574	0.001574	Pragmatic language
<i>KIAA0319</i>	0.01414	0.01301	0.04021	0.00289	0.00674	0.001015	0.004646	0.00076	0.00076	Pragmatic language
MinP	0.003648	0.000648	0.001037	0.002598	0.000177	0.000582	0.000244	0.00076	0.000177	Receptive language
MinP gene	<i>CNTNAP2</i>	<i>CNTNAP2</i>	<i>CMIP</i>	<i>CNTNAP2</i>	<i>CNTNAP2</i>	<i>CMIP</i>	<i>CNTNAP2</i>	<i>KIAA0319</i>	<i>CNTNAP2</i>	

SNPs with a nominal P-value of less than 9.9×10^{-4} are highlighted in bold. Minimum P values are given for all traits and genes.

No single SNP was significantly associated after multiple-testing corrections.

Table 3

Polygenic prediction of measures and outcomes in replication cohort

Consistency of polygenic profile score (base trait same as target trait)

Trait modelled	Best-fit threshold P-value	No. SNPs in best-fit model	Proportion of trait variance explained by polygenic score (R^2)	Nominal P-value ^a	Benjamini-Hochberg adjusted P-value ^b
Vocabulary 15months	0.09	32	0.106%	0.125	0.432
Vocabulary 24months	0.03	13	0.023%	0.480	0.549
Receptive language 15months	0.50	111	0.034%	0.385	0.513
Grammar 24months	0.16	47	0.089%	0.162	0.432
Receptive language 8years	0.03	19	0.061%	0.239	0.478
Expressive language 8years	0.23	65	0.182%	0.042	0.336
Nonword repetition 8years	0.09	42	0.008%	0.666	0.666
Pragmatic language 9years	0.01	9	0.056%	0.300	0.480

Overlaps in genetic effects (ability of expressive language polygenic profile to predict language and psychosocial outcomes at age 11)

Base Trait	Target Trait	Best-fit threshold P-value	No. SNPs in best-fit model	Proportion of trait variance explained by polygenic score (R^2)	Nominal P-value ^a	Benjamini-Hochberg adjusted P-value ^b
Expressive language 8years	DLD status	0.08	30	0.028%	0.419	0.628
Expressive language 8years	Emotional problems 11years	0.50	123	0.085%	0.210	0.420
Expressive language 8years	Peer problems 11years	0.06	27	0.428%	0.006	0.036
Expressive language 8years	Conduct problems 11years	0.01	9	0.094%	0.189	0.420
Expressive language 8years	Hyperactivity 11years	0.40	97	0.015%	0.601	0.666
Expressive language 8years	Prosociality 11years	0.02	17	0.010%	0.666	0.666

^aModels with nominal P-values less than 0.05 are highlighted in bold.^bBold results reached a significant level of association following a Benjamini-Hochberg adjustment at a False Discovery Rate of 0.05.

Figures

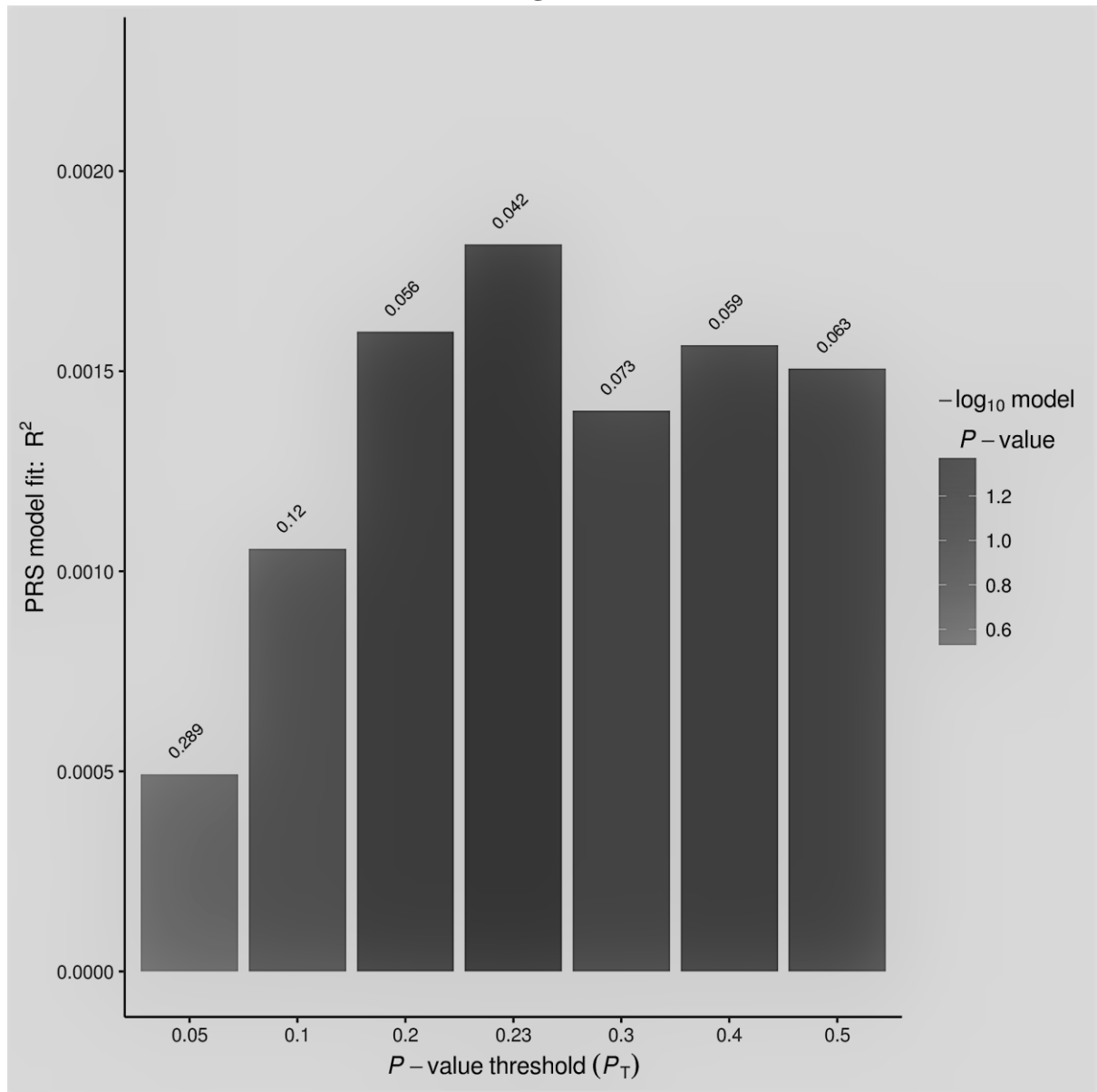


Figure 1. Best-fit model of genetic effects upon expressive language 8 years of age. The best-fit was found at a P-threshold of 0.23 and explains 0.18% of trait variance (R^2) in the replication sample ($P=0.042$).

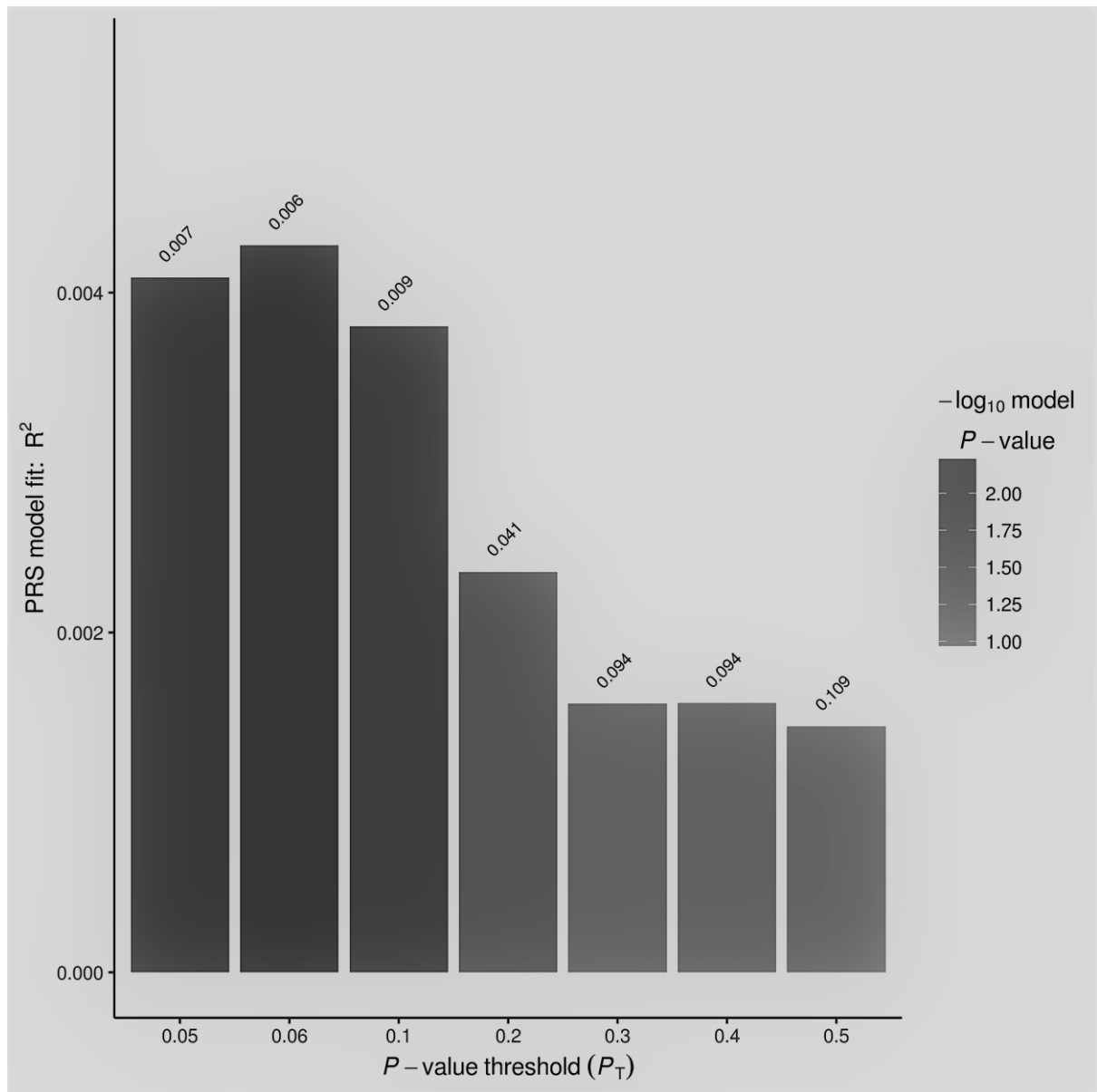


Figure 2. Best-fit model of genetic overlaps between expressive language at 8 years of age and peer problems.

The best-fit was found at a P-threshold of 0.06 and explains 0.43% of variance (R^2) in the replication sample ($P=0.006$).

Appendices

Table A1

Group level comparisons for all variables of interest

	n	Overall Range	Overall Mean (SD)	Without DLD Mean (SD)	With DLD Mean (SD)	Mean Difference [95% CI]	Test Statistics	Effect Size
Language and Communication Measures								
Vocabulary 15months	5259	0-268	88.36 (43.23)	89.37 (43.32)	73.43 (39.03)	15.94 [11.18, 20.71]	t(5257)=6.56***	0.37
Vocabulary 24months	5205	0-246	158.66 (53.16)	161.57 (51.88)	116.11 (53.34)	45.46 [39.69, 51.23]	t(5203)=15.44***	0.87
Receptive 15months	5263	0-12	9.18 (2.41)	9.20 (2.40)	8.90 (2.50)	0.30 [0.03, 0.57]	t(5261)=2.20*	0.12
Grammar 24months	5205	0-8	3.62 (2.55)	3.74 (2.53)	1.84 (2.05)	1.90 [1.63, 2.18]	t(5203)=13.42***	0.76
Receptive language 8years	5429	2-15	7.61 (1.87)	7.73 (1.80)	5.86 (1.99)	1.87 [1.68, 2.07]	t(5427)=18.62***	1.03
Expressive language 8years	5416	0-10	7.65 (1.68)	7.74 (1.61)	6.26 (2.04)	1.48 [1.30, 1.66]	t(5414)=16.22***	0.90
Non-word repetition 8years	5424	0-12	7.41 (2.42)	7.64 (2.27)	4.11 (2.13)	3.53 [3.28, 3.78]	t(5422)=28.03***	1.56
Pragmatic language 9years	4464	98- 162	151.83 (6.62)	152.43 (5.87)	144.12 (10.03)	8.32 [7.61, 9.03]	t(4462)=22.94***	1.33
Psychosocial Outcomes								
Emotional problems 11years	4308	0-10	1.36 (1.64)	1.34 (1.64)	1.65 (1.744)	-0.31 [-0.51, -0.11]	t(4306)=-3.05**	-0.19
Peer problems 11years	4157	0-9	0.95 (1.39)	0.92 (1.36)	1.47 (1.70)	-0.55 [-0.72, -0.38]	t(4155)=-6.37***	-0.40
Conduct problems 11years	4290	0-9	1.11 (1.33)	1.08 (1.31)	1.56 (1.56)	-0.49 [-0.65, -0.33]	t(4288)=-6.02***	-0.37
Hyperactivity 11years	4283	0-10	2.56 (2.13)	2.48 (2.08)	3.76 (2.50)	-1.28 [-1.54, -1.03]	t(4281)=-9.83***	-0.61
Prosociality 11years	4305	0-10	8.40 (1.62)	8.41 (1.62)	8.20 (1.69)	0.22 [0.02, 0.41]	t(4303)=2.18*	0.13

* p<.05, ** p<.01, *** p<.001.

Table A2.

Summary of SNPs by Gene

Gene	Number of SNPs in analysis	ALSPAC co-ordinates (hg38)
CMIP	172	chr16:81445241-81709799
ATP2C2	158	chr16:84368615-84463732
DCDC2	100	chr6:24174729-24357750
KIAA0319	52	chr6:24541241-24645764
FOXP2	27	chr7:114426511-114693772
CNTNAP2	720	chr7:146116876-148415616

Table A3.

SNPs in analyses which reached nominal level of significance ($P=10^{-4}$)

Chr	Position (hg38)	SNP	Gene	Allele1	included in best fit exp_8y profile ($P=0.23$)?	Vocabulary 15months		Vocabulary 24months		Receptive language 15months		Grammar 24months	
						BETA	P	BETA	P	BETA	P	BETA	P
6	24,314,672	rs9379651	DCDC2	A		-1.351	0.5093	-6.109	0.01607	-0.1064	0.336	-0.1663	0.1661
6	24,583,953	rs6902039	KIAA0319	C		1.16	0.3863	-0.8389	0.6099	0.149	0.04021	-0.04688	0.5469
7	146,727,562	rs34084487	CNTNAP2	C		5.07	0.07184	7.158	0.04169	0.2534	0.09833	0.5021	0.002598
7	147,487,190	rs6464815	CNTNAP2	T		0.6864	0.7096	-3.496	0.123	0.03331	0.7392	-0.1422	0.1853
7	147,572,753	rs12539980	CNTNAP2	C		4.956	0.003648	7.165	0.000648	0.2057	0.02591	0.2086	0.03608
7	147,818,988	rs17824995	CNTNAP2	G		1.414	0.5182	3.175	0.2367	0.008313	0.9441	0.08794	0.4891
7	147,889,090	rs10488350	CNTNAP2	C		1.838	0.3275	3.33	0.1491	-0.0247	0.8085	0.1546	0.1573
7	148,026,296	rs12533565	CNTNAP2	A		0.5879	0.6871	0.1989	0.9118	-0.02252	0.7765	0.000354	0.9967
7	148,038,344	rs17545705	CNTNAP2	T		1.214	0.4798	0.9511	0.6518	-0.04772	0.6083	0.01602	0.8724
7	148,361,425	rs9648690	CNTNAP2	G		-0.7228	0.5916	-2.727	0.09997	-0.1181	0.1057	-0.03607	0.6467
16	81,523,885	rs4889330	CMIP	T		-4.449	0.0944	-4.453	0.1764	-0.477	0.001037	0.01413	0.9278
16	81,657,207	rs8061723	CMIP	G	y	0.6783	0.7325	-0.12	0.961	0.07432	0.4901	0.03503	0.7627
16	84,373,185	rs12716749	ATP2C2	G		2.221	0.08795	0.01276	0.9937	0.1107	0.1165	0.1219	0.1079
16	84,375,135	rs4782938	ATP2C2	C		0.3783	0.7795	-2.336	0.1598	0.02716	0.7109	-0.00841	0.9149

Minimum p-value for each outcome measures is highlighted in bold. Tests which reach nominal p-value of 10^{-4} are highlighted in grey.

Table A3. (continued)

SNPs in analyses which reached nominal level of significance ($P=10^{-4}$)

Chr	Position (hg38)	SNP	Gene	Allele1	included in best fit exp_8y profile ($P=0.23$)?	Receptive language 8years		Expressive language 8years		Nonword repetition 8years		Pragmatic language 9years	
						BETA	P	BETA	P	BETA	P	BETA	P
6	24,314,672	rs9379651	DCDC2	A		-0.3094	0.000295	-0.05658	0.4727	-0.05744	0.6122	-0.2796	0.3931
6	24,583,953	rs6902039	KIAA0319	C		-0.07661	0.1682	0.02487	0.627	-0.1593	0.03062	-0.7102	0.00076
7	146,727,562	rs34084487	CNTNAP2	C		-0.1638	0.1674	0.1272	0.2427	0.02013	0.8977	-0.4656	0.3105
7	147,487,190	rs6464815	CNTNAP2	T		-0.1425	0.06321	-0.1227	0.08263	-0.1671	0.1005	-0.9668	0.000922
7	147,572,753	rs12539980	CNTNAP2	C		-0.03939	0.5783	-0.00515	0.9373	-0.00171	0.9854	-0.03829	0.8876
7	147,818,988	rs17824995	CNTNAP2	G		-0.07544	0.4059	-0.00309	0.9706	0.4062	0.000717	-0.0116	0.9733
7	147,889,090	rs10488350	CNTNAP2	C		-0.04349	0.5775	-0.02074	0.7733	0.3762	0.000275	-0.04313	0.8841
7	148,026,296	rs12533565	CNTNAP2	A		-0.02775	0.6457	-0.04145	0.4591	0.2951	0.000244	0.002313	0.9921
7	148,038,344	rs17545705	CNTNAP2	T		-0.05244	0.4618	-0.05376	0.4124	0.3128	0.000923	-0.1206	0.659
7	148,361,425	rs9648690	CNTNAP2	G		-0.2103	0.000177	-0.03933	0.4471	-0.158	0.03438	-0.2025	0.3485
16	81,523,885	rs4889330	CMIP	T		-0.02818	0.8012	0.04406	0.6692	-0.07686	0.6047	-0.01377	0.9739
16	81,657,207	rs8061723	CMIP	G	y	-0.07487	0.366	-0.2621	0.000582	-0.1408	0.1998	0.08177	0.7936
16	84,373,185	rs12716749	ATP2C2	G		-0.0351	0.518	-0.1695	0.000669	0.1498	0.0369	-0.1631	0.4294
16	84,375,135	rs4782938	ATP2C2	C		-0.01576	0.7792	-0.1766	0.000662	0.06389	0.3915	-0.3851	0.07137

Minimum p-value for each outcome measures is highlighted in bold. Tests which reach nominal p-value of 10^{-4} are highlighted in grey.